

APPENDIX B

to declaration by Tetsuya Gatanaga, Ph.D.

Data from Clinical Trial, Protocol AIT-PAN-20

Survival
 CYTOIMPLANT

Site/Patient Number	Patient Initials	Age (years)	Sex	Baseline Disease Stage	Date of Randomization	Date of Death	Survival Days (as of 10-Nov-99)	Status (as of 10-Nov-99)
01-002	MSM	67	F	Stage IV, T4 N1 Mx	27-Jan-99	24-May-99	117	Deceased
01-004	HGK	73	M	Stage IV, T4 N0 M0	09-Mar-99	24-Aug-99	168	Deceased
01-005	MAW	67	F	Stage IV, T4 N0 Mx	31-Mar-99	N/A	224+	Off Study
01-006	LCT	64	F	Stage IV, T4 N1 Mx	01-Apr-99	14-Jul-99	104	Deceased
01-009	EAR	58	F	Stage IV, T4 N1 M1	20-Apr-99	N/A	204+	Off Study
01-010	DLW	78	M	Stage III, T3 N1 Mx	30-Jul-99	N/A	103+	On Study
02-052	A-P	65	F	Stage IV, T4 N0 M0	25-Feb-99	22-May-99	86	Deceased
02-053	S-M	33	F	Stage IV, T4 Nx M1	29-Mar-99	21-Jun-99	84	Deceased
02-054	CEM	58	M	Stage IV, T4 N1 M1	27-Apr-99	26-Jun-99	60	Deceased
03-101	REW	54	M	Stage III, T3 N1 M0	08-Feb-99	N/A	275+	On Study
03-102	DLD	53	M	Stage IV, T4 N1 M0	18-Mar-99	N/A	237+	Off Study
03-104	VFB	85	F	Stage IV, T4 N0 M0	04-May-99	N/A	190+	On Study
03-105	AMS	79	M	Stage IV, T4 N1 M1	14-Jun-99	14-Oct-99	122	Deceased
03-108	WLG	57	M	Stage IV, T4 N0 M1	07-Sep-99	N/A	64+	On Study
04-151	M-B	78	F	Stage II, T3 N0 M0	11-Mar-99	20-May-99	70	Deceased
04-152	EMD	71	F	Stage III, T3 N1 M0	09-Jun-99	N/A	154+	Off Study
05-203	BFB	65	F	Stage IV, T4 N1 Mx	07-Jul-99	01-Aug-99	25	Deceased
05-204	PAQ	70	M	Stage IV, T4 N0 M1	19-Jul-99	N/A	114+	On Study
05-206	RES	78	M	Stage II, T3 N0 M0	25-Aug-99	N/A	77+	On Study
05-207	RJW	74	M	Stage III, T2 N1 M0	14-Sep-99	N/A	57+	On Study
05-208	MJS	69	F	Stage II, T3 Nx Mx	18-Oct-99	N/A	23+	On Study
06-251	FJM	63	M	Stage III, T3 N1 M0	15-Oct-99	N/A	26+	On Study
08-351	P-F	63	F	Stage IV, T4 N1 Mx	12-Oct-99	N/A	29+	On Study

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Protocol ATT-PAN-201

Survival

Gemcitabine

Site/Patient Number	Patient Initials	Age (years)	Sex	Baseline Disease Stage	Date of Randomization	Date of Death	Survival Days (as of 10-Nov-99)	Status (as of 10-Nov-99)
01-001	CDR	81	M	Stage IV, T4 N1 M0	10-Dec-98	03-Nov-99	328	Deceased
01-003	WMH	55	F	Stage III, T3 N1 M0	15-Feb-99	N/A	268+	Off Study
01-007	REH	53	M	Stage IV, T4 N0 M0	06-Apr-99	N/A	218+	On Study
01-008	ADC	59	F	Stage IV, T4 N1 M1	13-Apr-99	17-Aug-99	126	Deceased
02-051	CHS	75	M	Stage IV, T4 N0 M1	11-Feb-99	N/A	272+	Off Study
02-055	LDR	50	M	Stage IV, T4 N1 M0	10-Sep-99	N/A	61+	On Study
03-103	EFK	75	F	Stage IV, T4 N0 M0	23-Apr-99	N/A	201+	On Study
03-106	R-M	72	M	Stage IV, T3 N0 M1	09-Jul-99	N/A	124+	On Study
03-107	CGD	60	F	Stage II, T3 N0 M0	19-Jul-99	N/A	114+	Off Study
05-201	JLB	79	F	Stage III, T1 N1 M0	09-Jun-99	N/A	154+	Off Study
05-202	D-S	59	M	Stage IV, T3 N0 M1	25-Jun-99	N/A	138+	On Study
05-205	EMS	72	F	Stage IV, T4 N0 M0	04-Aug-99	N/A	98+	On Study
07-301	RIW	71	F	Stage IV, T2 N1 M1	18-Oct-99	N/A	23+	On Study

Survival Days = Date of Death (or 10-Nov-99) - Randomization Date

N/A = Not Applicable (death not reported for patient as of 10-Nov-99)

Patients 003 and 107 withdrew from the study upon randomization to Gemcitabine.

Time to Treatment Failure
CYTOIMPLANT

Site/Patient Number	Patient Initials	Age (years)	Sex	Baseline Disease Stage	Date of Randomization	Date of Treatment Failure	Time to Treatment Failure (days)	Reason for Treatment Failure
01-002	MSM	67	F	Stage IV, T4 N1 M0	27-Jan-99	27-Apr-99	90	Progressive Disease
004	HGK	73	M	Stage IV, T4 N0 M0	09-Mar-99	02-Jun-99	85	Progressive Disease
01-005	MAW	67	F	Stage IV, T4 N0 M0	31-Mar-99	07-Jul-99	98	Progressive Disease
01-006	LCT	64	F	Stage IV, T4 N1 M0	01-Apr-99	19-May-99	48	Progressive Disease
01-009	EAR	58	F	Stage IV, T4 N1 M1	20-Apr-99	21-Jul-99	92	Progressive Disease
01-010	DLW	78	M	Stage III, T3 N1 M0	30-Jul-99	25-Oct-99	87	Progressive Disease
02-052	A-P	65	F	Stage IV, T4 N0 M0	25-Feb-99	22-May-99	86	Death
02-053	S-M	33	F	Stage IV, T4 N0 M1	29-Mar-99	21-Jun-99	84	Death
02-054	CEM	58	M	Stage IV, T4 N1 M1	27-Apr-99	27-May-99	30	Progressive Disease
03-101	REW	54	M	Stage III, T3 N1 M0	08-Feb-99	28-Sep-99	232	Progressive Disease
03-102	DLD	53	M	Stage IV, T4 N1 M0	18-Mar-99	25-Jun-99	99	Progressive Disease
03-104	VFB	85	F	Stage IV, T4 N0 M0	04-May-99	25-Oct-99	174	Progressive Disease
03-105	AMS	79	M	Stage IV, T4 N1 M1	14-Jun-99	16-Jul-99	32	Progressive Disease
108	WLG	57	M	Stage IV, T4 N0 M1	07-Sep-99	N/A	N/A	
04-151	M-B	78	F	Stage II, T3 N0 M0	11-Mar-99	20-May-99	70	Death
04-152	EMD	71	F	Stage III, T3 N1 M0	09-Jun-99	10-Aug-99	62	Progressive Disease
05-203	BFB	65	F	Stage IV, T4 N1 M0	07-Jul-99	01-Aug-99	25	Death
05-204	PAQ	70	M	Stage IV, T4 N0 M1	19-Jul-99	N/A	N/A	
05-206	RES	78	M	Stage II, T3 N0 M0	25-Aug-99	N/A	N/A	
05-207	RJW	74	M	Stage III, T2 N1 M0	14-Sep-99	N/A	N/A	
05-208	MJS	69	F	Stage II, T3 N0 M0	18-Oct-99	N/A	N/A	
06-251	FJM	63	M	Stage III, T3 N1 M0	15-Oct-99	N/A	N/A	
08-351	P-F	63	F	Stage IV, T4 N1 M0	12-Oct-99	N/A	N/A	

Protocol ALT-PAN-201
Time to Treatment Failure
Gemcitabine

Site/Patient Number	Patient Initials	Age (years)	Sex	Baseline Disease Stage	Date of Randomization	Date of Treatment Failure	Time to Treatment Failure (days)	Reason for Treatment Failure
01-001	CDR	81	M	Stage IV, T4 N1 M0	10-Dec-98	07-Apr-99	118	Progressive Disease
01-003	WMH	55	F	Stage II, T3 N1 M0	15-Feb-99	23-Feb-99	8	Patient withdrew consent
01-007	REH	53	M	Stage IV, T4 N0 MX	06-Apr-99	11-Jun-99	66	Progressive Disease
01-008	ADC	59	F	Stage IV, T4 N1 M1	13-Apr-99	17-Aug-99	126	Death
02-051	CHS	75	M	Stage IV, T4 N1 M1	11-Feb-99	21-Sep-99	222	Progressive Disease
02-055	LDR	50	M	Stage IV, T4 N1 M0	10-Sep-99	N/A	N/A	
03-103	EFK	75	F	Stage IV, T4 N0 MX	23-Apr-99	N/A	N/A	
03-106	R-M	72	M	Stage IV, T3 N0 M1	09-Jul-99	N/A	N/A	
03-107	CGD	60	F	Stage II, T3 N0 M0	19-Jul-99	19-Jul-99	0	Patient withdrew consent
05-201	JLB	79	F	Stage II, T1 N1 M0	09-Jun-99	29-Jul-99	50	Patient withdrew consent
05-202	D-S	59	M	Stage IV, T3 N0 M1	25-Jun-99	N/A	N/A	
05-205	EMS	72	F	Stage IV, T4 N0 M0	04-Aug-99	N/A	N/A	
201	RIW	71	F	Stage IV, T2 N1 M1	18-Oct-99	N/A	N/A	

Time to Treatment Failure = Date of Treatment Failure - Date of Randomization

N/A = Not Applicable (no Treatment Failure as of 10-Nov-99)

Patients 003 and 107 withdrew from the study upon randomization to Gemcitabine.

Patient 201 refused to continue therapy.

Adverse Events Possibly/Probably Related to Injection Procedure and/or Study Drug
AE/SAE Listings

Randomized Treatment: Cytoimplant

Investigator	Pat Init	Date of Rard	Date of 1st Treatment	Date of Last Treatment	Date of Onset	Adverse Event	SAE Number	Resolut.	Severity	Relationship to:	
										Injection Procedure	Study Drug
1/HAMES, ROBERT	2	MSH	27JAN1999	02FEB1999	-	15APR1999 VOMITING	1999-WCO346	18MAY1999	Sev	N/A	Poss
						ABDOMINAL PAIN	1999-WCO346	18MAY1999	Sev	N/A	Poss
	10	DLW	30JUL1999	25AUG1999	-	25AUG1999 ABDOMINAL PAIN	26AUG1999		Mod	Poss	Poss
2/ERICKSON, RICHARD	52	A-P	25FEB1999	08MAR1999	-	09MAR1999 CHILLS	22MAY1999		Mild	Poss	Poss
	53	S-H	29MAR1999	06APR1999	-	15APR1999 FEVER	29APR1999		Mild	Prob	Prob
	54	CEH	27APR1999	04MAY1999	-	04MAY1999 FEVER	26JUN1999		Mild	Poss	Poss
3/KOZAREK, RICHARD	101	REW	08FEB1999	17FEB1999	21JUL1999	17FEB1999 HOT FLASH	17FEB1999		Mild	Prob	Prob
						LOW BACK PAIN	22FEB1999		Mod	Prob	Prob
						OBSTIPATION	23FEB1999		Mod	Poss	Poss
						DARK URINE	22FEB1999		Mod	Prob	Prob
						UPPER GIRTH PAIN	05APR1999		Mod	Prob	Prob
						DECREASED APPETITE	22JUL1999		Mild	Prob	Prob
						20FEB1999 CONSTIPATION	22FEB1999		Mod	Poss	Poss
						23FEB1999 PANCREATIC	05APR1999		Mod	Prob	Prob
						INFILAMMATION					
						BACK PAIN	03MAY1999		Sev	Poss	Poss
						04MAR1999 BACK PAIN	22JUL1999		Mod	Poss	Poss
						24MAR1999 NAUSEA	06APR1999		Sev	Prob	Prob
						VOMITING	06APR1999		Mod	Prob	Prob
						30MAR1999 EPIGASTRIC BURNING	06APR1999		Sev	Prob	Prob
						30MAR1999 GASTRIC OUTLET	1999-WCO263		Sev	Prob	Prob
						OBSSTRUCTION	06APR1999				
						104 VFB	10MAY1999		Mod	Prob	Prob
						04MAY1999 20SEP1999	12MAY1999		Mod	Unlk	Unlk
						10MAY1999 10MAY1999	24MAY1999		Mod	Prob	Prob
						EPIGASTRIC BURNING					
						EPIGASTRIC					
						DISCOMFORT					

Date of Last Treatment (Cytoimplant patients) = - ==> patient not treated with second Cytoimplant as of data cut-off.
 Date of Resolution = - ==> if date is not given, event is ongoing or date of resolution is unavailable.
 Special Missing values: N = Not Applicable; D = Not Done; U = Unknown
 All Adverse Events associated with an SAE number have been reported as a Serious Adverse Event only if YES is noted in the Serious column.
 More than one adverse event may have been reported with an SAE. These events are the signs and symptoms of the diagnosis.

AE/SAE Listings

Investigator	Pat Init	Date of Rand	Date of	Date of	Adverse Event	SAE Number	Date of Resolut.	Severity	Relationship to:		
			1st Treatment	Last Treatment Onset					Injection Procedure	Study Drug	
3/KOZAREK, RICHARD	104	VFB	04MAY1999	10MAY1999	20SEP1999	11MAY1999	FEVER, 37.7 C NAUSEA VOMITING CHILLS DIARRHEA ABDOMINAL PAIN DEHYDRATION	12MAY1999 1999-WC0391 1999-WC0391	Mod Sev Mod Mod Mod Mod Mod	Prob Prob Prob Prob Prob Prob Prob	Prob Prob Prob Prob Prob Prob Prob
105 AMS	14JUN1999	23JUN1999	18AUG1999	23JUN1999	19AUG1999	11MAY1999	NAUSEA/VOMITING RIGORS	24JUN1999 19AUG1999	Mod Mod	Poss Prob Prob	Poss Prob Prob
108 MLC	07SEP1999	21SEP1999	-	23SEP1999	FEVER	23MAR1999	INCREASE IN TEMPERATURE	24SEP1999	Mod	Poss	Poss
4/GRESS, FRANK	151 M-B	11MAR1999	22MAR1999	-	23MAR1999	15JUN1999	GASSINESS NAUSEA VOMITING ABDOMINAL DISCOMFORT	30MAR1999	Mod	Poss	Unlk
5/NGUYEN, CUONG	204 PAB	09JUN1999	16JUN1999	-	16JUN1999	20JUL1999	FEVER DIARRHEA	17AUG1999 02AUG1999	Mod Mod	Poss Poss Poss Poss	Unlk Unlk Unlk Unlk
208 MJS	18OCT1999	28OCT1999	-	28OCT1999	PANCREATITIS FECAL INCONTINENCE	1999-WC0742	05NOV1999	Mod Mod Mod Mod	Poss Poss Poss Prob	Poss Poss Poss Unlk	
8/FAIGEL, DOUGLAS O.	351 P-F	12OCT1999	19OCT1999	-	19OCT1999	NAUSEA	19OCT1999 29OCT1999	Mod Mod	Poss Poss	Poss Poss	

Date of Last treatment (Cytoimplant patients) = - ==> Patient not treated with second Cytoimplant as of data cut-off.
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AE/SAE Listings
Related to Injection Procedure and/or Study Drug

Randomized Treatment: Cytoimplant

Investigator	Pat Init	Pat Rand	Date of 1st Treatment	Date of Last Treatment	Date of Onset	Adverse Event	SAE Number	Relationship to:		
								Injection Procedure	Study Drug	Relationship to:
8/FAIGEL, DOUGLAS O.	351 P-F	120C1999	190C1999	-	190C1999	ABDOMINAL TENDERNESS	290C1999	Mild	Poss	Poss

Date of Last treatment (Cytoimplant patients) = - ==> Patient not treated with second Cytoimplant as of data cut-off.
Date of Resolution = - ==> if date is not given, event is ongoing or date of resolution is unavailable.
Special Missing values: N = Not Applicable; D = Not Done; A = Not Available; U = Unknown
All Adverse Events associated with an SAE number have been reported as a Serious Adverse Event only if YES is noted in the Serious column.
More than one adverse event may have been reported with an SAE. These events are the signs and symptoms of the diagnosis.

Randomized Treatment: Gemcitabine

Investigator	Pat Init	Date of 1st Rand	Date of Treatment	Date of Last Treatment	Date of Onset	Adverse Event	SAE Number	Date of Resolut.	Severity	Relationship to:	
										Injection Procedure	Study Drug
1/HAWES, ROBERT	1 CDR	10DEC1998	18DEC1998	22DEC1998	31DEC1998	MOUTH SORES	08JAN1999	Mild	N/A	Prob	Prob
						PLATELET COUNT LOW (33K)	08JAN1999	Mod	N/A	Prob	Prob
						PLATELET COUNT LOW (44K)	27JAN1999	Mod	N/A	Prob	Prob
						12MAR1999 LOW PLAT. COUNT (33)	19MAR1999	Mod	N/A	Prob	Prob
						16APR1999 HEMOGLOBIN 8.6	16APR1999	Mod	N/A	Prob	Prob
						23APR1999 LOW DECREASED PLATELET COUNT (60K)	30APR1999	Mod	N/A	Prob	Prob
						28MAY1999 LOW DECREASED PLAT. COUNT (39K)	04JUN1999	Mod	N/A	Prob	Prob
						02JUL1999 LOW PLAT. COUNT (28K)	09JUL1999	Mod	N/A	Prob	Prob
						LOW WBC (1.6)	09JUL1999	Mod	N/A	Prob	Prob
						LOW PLATELET COUNT 28	09JUL1999	Mod	N/A	Prob	Prob
						23JUL1999 WBC 3.9 LOW	13AUG1999	Mild	N/A	Prob	Prob
						30JUL1999 WBC 2.4 LOW	13AUG1999	Mod	N/A	Prob	Prob
						03AUG1999 LOW PLAT. COUNT 31	13AUG1999	Mod	N/A	Prob	Prob
						WBC 1.6 LOW	13AUG1999	Mod	N/A	Prob	Prob
						03SEP1999 WBC 3.0 LOW	17SEP1999	Mild	N/A	Prob	Prob
						10SEP1999 LOW PLAT. COUNT 36	17SEP1999	Mod	N/A	Prob	Prob
						WBC 1.7 LOW	17SEP1999	Mod	N/A	Prob	Prob
						01OCT1999 LOW HEMOGLOBIN 8.2	15OCT1999	Mild	N/A	Prob	Prob
						08OCT1999 LOW HEMOGLOBIN 8.3	15OCT1999	Mild	N/A	Prob	Prob
						WBC 3.2 LOW	22OCT1999	Mild	N/A	Prob	Prob
						15OCT1999 LOW PLAT. COUNT 16K	15OCT1999	Sev	N/A	Prob	Prob
						LOW HEMOGLOBIN 8.4	15OCT1999	Mild	N/A	Prob	Prob

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All Adverse Events associated with an SAE number have been reported as a Serious Adverse Event only if YES is noted in the Serious column.

More than one adverse event may have been reported with an SAE. These events are the signs and symptoms of the diagnosis.

Randomized Treatment: Gemcitabine

Investigator	Pat Init	Date of Rand	Date of Treatment	1st Treatment	Last Treatment	Date of Onset	Date of Adverse Event	SAE Number	Date of Resolut.	Severity	Injection Procedure	Study Drug
1/HAWES, ROBERT	1 CDR 7 REH	10DEC1998 06APR1999	18DEC1998 09APR1999	22OCT1999 16AUG1999	15OCT1999 20MAY1999	WBC 1.3 HEMOGLOBIN 6.0	LOW DECREASED DECREASED	22OCT1999 20MAY1999	Mod Mod	N/A N/A	Prob Prob	
						HEMATOCRIT 20	DECREASED	20MAY1999	Mod	N/A	Prob	
						06JUL1999	DECREASED	23JUL1999	Mod	N/A	Prob	
						HEMOGLOBIN (7.7)	DECREASED	23JUL1999	Mod	N/A	Prob	
						HEMATOCRIT (23)	DECREASED	23JUL1999	Mod	N/A	Prob	
						LOW PLATELETS 71K	LOW PLATELETS 60K	19MAY1999 19MAY1999	Mod Mod	N/A N/A	Prob Prob	
						ANC, LOW 880	ANC, LOW 880	19MAY1999 03JUN1999	Mod Mod	N/A N/A	Prob Prob	
						LOW HEMOGLOBIN	LOW HEMOGLOBIN	24JUN1999	Mod	N/A	Prob	
						LOW PLATELET 82	LOW PLATELET 82	21JUL1999	Mod	N/A	Prob	
						07JUL1999	LOW PLATELET 95	28JUL1999	Mod	N/A	Prob	
						ANC, LOW 700						
2/ERICKSON, RICHARD	51 CHS	11FEB1999	16FEB1999	14SEP1999	18FEB1999	EMESIS		16MAR1999	Mod	N/A	Pass	
		19FEB1999	23FEB1999	DEPRESSION CONSTIPATION					Mod	N/A	Pass	
				ACID REFLUX					Mod	N/A	Pass	
				06APR1999	SHORNESS OF BREATH				Mod	N/A	Pass	
				06APR1999	DRY MOUTH				Mod	N/A	Pass	
				21SEP1999	SHORNESS OF BREATH				Mod	N/A	Pass	
				29SEP1999	JOINT STIFFNESS				Mod	N/A	Pass	
				29SEP1999	HEADACHE				Mod	N/A	Pass	
				29SEP1999	HOT FLASHERS				Mod	N/A	Pass	

Adverse Events Possibly/Probably Related to Injection Procedure and/or Study Drug

Randomized Treatment: Gemcitabine

Date: 15DEC1999
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Investigator	Pat Init	Date of Rand	Date of Treatment	Date of Treatment	Date of Onset	Adverse Event	SAE Number	Date of Resolut.	Severity	Relationship to:	
										Injection Procedure	study Drug
2/ERICKSON, RICHARD	55 LDR	10SEP1999	14SEP1999	21OCT1999	29SEP1999	CHILLS (L) SIDE CHEST PAIN		29SEP1999	Mod	N/A	Poss
					06OCT1999	NAUSEA		29SEP1999	Mod	N/A	Poss
					12OCT1999	WEAKNESS			Mod	N/A	Poss
						EPIGASTRIC PAIN			Mod	N/A	Poss
3/KOZAREK, RICHARD	103 EFK	23APR1999	30APR1999	17SEP1999	03MAY1999	RASH ON CHEST		06MAY1999	Mod	N/A	Poss
					13MAY1999	NEUTROPEMIA		20MAY1999	Mod	N/A	Prob
					27MAY1999	THROMBOCYTOPENIA		20MAY1999	Mod	N/A	Prob
					03JUN1999	NEUTROPEMIA		03JUN1999	Mod	N/A	Prob
					03JUN1999	NEUTROPEMIA		09JUN1999	Mod	N/A	Prob
					12JUL1999	MOUTH SORES		26JUL1999	Mod	N/A	Prob
					26JUL1999	VOMITING		26JUL1999	Mod	N/A	Prob
					26JUL1999	NAUSEA		26JUL1999	Mod	N/A	Prob
					02SEP1999	INCREASED FATIGUE FOLLOWING		30SEP1999	Mod	N/A	Prob
					03SEP1999	CHEMOTHERAPY				N/A	
					03SEP1999	NUMBNESS IN FEET				N/A	
					18SEP1999	NAUSEA				N/A	
					18SEP1999	LEG WEAKNESS				N/A	
					22SEP1999	VOMITING				N/A	
						DEPRESSION				N/A	
						FEVER				N/A	
						04AUG1999				N/A	
						04AUG1999	NAUSEA			N/A	
						11AUG1999	FATIGUE			N/A	
						24AUG1999	ANEMIA			N/A	
						OBSEP1999	I.V. INFILTRATION			N/A	
						25AUG1999				N/A	
						08SEP1999				N/A	

Date of Last Treatment (Cyroimplant patients) = - ==> Patient not treated with second Cyroimplant as of date cut-off.

Date of Resolution = ==> if date is not given, event is ongoing or, date of resolution is unavailable.

Special Missing values: N = Not Applicable; D = Not Done; A = Not Available; U = Unknown

All Adverse Events associated with an SAE number have been reported as a Serious Adverse Event only if YES is noted in the Serious column.

More than one adverse event may have been reported with an SAE. These events are the signs and symptoms of the diagnosis.

Randomized Treatment: Gemcitabine

Date: 15DEC1999
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Investigator	Pat Init	Date of Random	Date of 1st Treatment	Date of Last Treatment	Date of Onset	Adverse Event	SAE Number	Date of Resolut.	Severity	Relationship to:	
										Injection Procedure	Study Drug
3/KOZAREK, RICHARD	106 R-M	09JUL1999	14JUL1999	200CT1999	200CT1999	NEUTROPENIA	03NOV1999	Mild	N/A	Prob	
5/NGUYEN, CUONG	201 JLB	09JUN1999	21JUN1999	13JUL1999	06JUL1999	THROMBOCYTOPENIA	13JUL1999	Mild	N/A	Prob	
					13JUL1999	ANEMIA	UUAUUAUUAU	Mild	N/A	Prob	
					15JUL1999	GALLBLADDER FLUID	UUAUAG1999	Mild	N/A	Poss	
						INFECTION +CULTURE		Mod	N/A	Poss	
						GRAM NEG RODS					
						ANEMIA					
205 EMS	06AUG1999	13AUG1999	18OCT1999	03SEP1999							

Date of Last Treatment (Cytomim patient) = - ==> Patient not treated with second Cytomim as of data cut-off.
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 Special Missing values: N = Not Applicable; D = Not Done; A = Not Available; U = Unknown
 All Adverse Events associated with an SAE number have been reported as a Serious Adverse Event only if YES is noted in the Serious column.
 More than one adverse event may have been reported with an SAE. These events are the signs and symptoms of the diagnosis.